General Describio References Comments Links General Features

### General information

Billianding ... LIP1 CANRU

P20261 

XP-002230351

Created : . . .

Rel. 17, 1-FEB-1991

Sequence update Rel. 27, 1-OCT-1993

Annotation update: Rel. 41, 15-JUN-2002

PD 01-02-150

### Description and origin of the Protein

Description:

Lipase 1 precursor (EC 3.1.1.3).

Gene name(s)

LIP1.

Organism source Taxonomy ?

Candida rugosa (Yeast) (Candida cylindracea).

Eukaryota; Fungi; Ascomycota; Saccharomycotina; Saccharomycetes; Saccharomycetales;

mitosporic Saccharomycetales; Candida.

NCBI TaxID

5481

THE ST

#### References

[1] Longhi, S., Fusetti, F., Grandori, R., Lotti, M., Vanoni, M., Alberghina, L., Cloning and nucleotide sequences of two lipase genes from Candida cylindracea. (1992) Biochim. Biophys. Acta 1131:227-232

Position SEQUENCE FROM N.A.

Comments STRAIN=ATCC 14830;

Medline 92305068

PubMed 1610906

[2] Kawaguchi, Y., Honda, H., Taniguchi-Morimura, J., Iwasaki, S., The codon CUG is read as serine in an asporogenic yeast Candida cylindracea. (1989) Nature 341:164-166

Position SEQUENCE OF 12-549 FROM N.A., AND PARTIAL SEQUENCE.

Comments STRAIN=ATCC 14830 / MS-5;

Medline 89384874

PubMed 2506450

[3] Grochulski, P., Li, Y., Schrag, J.D., Bouthillier, F., Smith, P., Harrison, D., Rubin, B., Cygler, M., Insights into interfacial activation from an open structure of Candida rugosa lipase. (1993) J. Biol. Chem. 268:12843-12847

Position X-RAY CRYSTALLOGRAPHY (2.06 ANGSTROMS).

Medline 93286131

PubMed 8509417

[4] Grochulski, P., Bouthillier, F., Kazlauskas, R.J., Serreqi, A.N., Schrag, J.D., Ziomek, E., Cygler, M., Analogs of reaction intermediates identify a unique substrate binding site in Candida rugosa lipase.

(1994) Biochemistry 33:3494-3500

Position X-RAY CRYSTALLOGRAPHY (2.05 ANGSTROMS).

Medline 94190867

PubMed 8142346

[5] Benjamin, S., Pandey, A.,

Candida rugosa lipases: molecular biology and versatility in biotechnology. (1998) Yeast 14:1069-1087

Position REVIEW.

Medline 98451816

#### PubMed 9778794

## Comments

CATALYTIC ACTIVITY

TRIACYLGLYCEROL + H(2)O = DIACYLGLYCEROL + A FATTY ACID

ANION.

多种人的 网络多克拉特 "我没有一个一

SIMILARITY

BELONGS TO THE TYPE-B CARBOXYLESTERASE/LIPASE FAMILY.

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#### Database cross-references

**EMBL** 

X64703;CAA45957.1;-.

X16712;CAA34684.1;-.

PIR

S05684;S05684.

S23448;S23448.

1CRL:31-JAN-94.

1LPM;20-APR-95.

1LPN;20-APR-95.

PDB

1LPO;20-APR-95.

1LPP:20-APR-95.

1LPS;08-MAR-95.

1TRH;31-JAN-94.

InterPro

IPR002018;CarbesteraseB.

IPR000379;Ser\_estrs\_site.

Pfam

PF00135;COesterase;1.

**PROSITE** 

PS00122; CARBOXYLESTERASE\_B\_1;1. PS00941;CARBOXYLESTERASE\_B\_2;1.

#### Keywords

Hydrolase; Lipid degradation; Signal; Glycoprotein; Multigene family; 3D-structure;

#### **Features**



Key	Beglio.	re End E	Length	Descrivo
SIGNAL	1	15	15	
CHAIN	16	549	534	LIPASE 1.
ACT_SITE	224	224	1	
ACT_SITE	356	356	1	
ACT_SITE	464	464	1	
DISULFID	75	112	38	
DISULFID	283	292	10	
CARBOHYD	329	329	1	N-LINKED (GLCNAC).
CARBOHYD	366	366	1	N-LINKED (GLCNAC).
VARIANT	398	398	1	G -> Q.

STRAND	18	20	3
TURN	22	23	2
STRAND	26	28	3
STRAND	30	31	2
STRAND	36	42	7
STRAND	44	44	1
HELIX	49	51	3
TURN	52	53	2
STRAND	58	58	1
TURN	64	65	2
STRAND	67	67	1
STRAND	69	69	1
STRAND	74	74	1
TURN	80	81	2
HELIX	88	97	10
TURN	98	98	1
HELIX	100	105	6
STRAND	108	108	1
STRAND	114	119	6
TURN	121	122	2
TURN	125	126	2
STRAND	129	135	7
TURN	139	141	3
TURN	146	147	2
HELIX	151	159	9
TURN	160	161	2
STRAND	165	169	5
HELIX	174	178	5
HELIX	182	187	6
TURN	188	188	1
TURN	190	191	2
HELIX	192	207	16
HELIX	208	211	4
TURN	212	212	1
STRAND	213	223	11
TURN	224	224	1
HELIX	225	235	11
HELIX	236	239	4
STRAND	242	243	2
TURN	244	245	2
STRAND	246	247	2
STRAND	251	255	5
TURN	265	266	2
HELIX	268	281	14

TURN	282	282	1
TURN	284	285	2
HELIX	289	295	7
HELIX	298	306	9
TURN	307	307	1
TURN	311	312	2
TURN	314	317	4
HELIX	334	339	6
TURN	340	341	2
STRAND	348	353	6
TURN	354	354	1
STRAND	355	355	1
TURN	356	357	2
HELIX	358	361	4
HELIX	362	364	3
TURN	365	366	2
HELIX	370	380	11
TURN	382	383	2
HELIX	386	395	10
HELIX	400	402	3
TURN	406	407	2
TURN	409	412	4
HELIX	418	429	12
TURN	430	430	1
HELIX	431	440	10
STRAND	446	451	6
TURN	453	456	4
TURN	458	460	3
STRAND	463	463	1
TURN	464	465	2
HELIX	466	472	7
TURN	473	473	1
TURN	477	478	2
HELIX	<b>4</b> 79	482	4
TURN	483	483	1
HELIX	484	492	9
HELIX	495	498	4
STRAND	507	507	1
TURN	510	511	2
STRAND	517	520	4
STRAND	525	525 528	
HELIX	534	541	8
HELIX	544	547	4

Sequence information

# Length: 549 aa, molecular weight: 58550 Da, CRC64 checksum: 27A40BD318757CE0

//

MELALALSLI						60
GSLDGOKFTS	YGPSCMQQNP	EGTYEENLPK	AALDLVMQSK	VFEAVSPSSE	DCLTINVVRP	120
PGTKAGANLP	VMLW1FGGGF	EVGGTSTFPP	AQMITKSIAM	GKPIIHVSVN	YRVSSWGFLA	180
GDEIKAEGSA	NAGLKDORLG	MQWVADNIAA	FGGDPTKVTI	FGESAGSMSV	MCHILWNDGD	240
NTYKGKPLFR	AGIMOSGAMV	PSDAVDGIYG	NEIFDLLASN	AGCGSASDKL	ACLRGVSSDT	300
LEDATNNTPG						360
GTSSLNVTTD	AOAREYFKOS	FVHASDAEID	TLMTAYPGDI	TQGSPFDTGI	LNALTPQFKR	420
ISAVLGDLGF	TLARRYFLNH	YTGGTKYSFL	SKQLSGLPVL	GTFHSNDIVF	QDYLLGSGSL	480
IYNNAFIAFA	TDLDPNTAGL	LVKWPEYTSS	SQSGNNLMMI	NALGLYTGKD	NFRTAGYDAL	540
FSNPPSFFV						549

General Description References Comments Links Kerwhits Features Sequence



# Title: EP02009616\_0010.PROT Application Date: 07-Feb-2003

>>SWALL:LIP1\_CANRU P20261 Lipase 1 precursor (EC 3.1.1.3 (549 aa) 01-Feb-1991 initn: 3466 init1: 3466 opt: 3468 Z-score: 3919.8 bits: 735.1 E(): 3.8e-210 Smith-Waterman score: 3468; 96.276% identity (96.276% ungapped) in 537 aa overlap (11-3.0 SMNSRGPAGRLGSVPTATLANGDTITGLNAIINEAFLGIPFAEPPVGNLRFKDPVPYS EP0200 SWALL: MELALALSLIASVAAAPTATLANGDTITGLNAIINEAFLGIPFAEPPVGNLRFKDPVPYS EP0200 GSLDGOKFTSYGPSCMQQNPEGTYEENLPKAALDLVMQSKVFEAVSPSSEDCLTINVVRP SWALL: GSLDGQKFTSYGPSCMQQNPEGTYEENLPKAALDLVMQSKVFEAVSPSSEDCLTINVVRP EP0200 PGTKAGANLPVMLWIFGGGFEVGGTSTFPPAQMITKSIAMGKPIIHVSVNYRVSSWGFLA SWALL: PGTKAGANLPVMLWIFGGGFEVGGTSTFPPAQMITKSIAMGKPIIHVSVNYRVSSWGFLA EP0200 GDEIKAEGSANAGLKDQRMGMQWVADNIAAFGGDPTKVTIFGESAGSMSVMCHILWNDGD SWALL: GDEIKAEGSANAGLKDQRLGMQWVADNIAAFGGDPTKVTIFGESAGSMSVMCHILWNDGD EP0200 NTYKGKPLFRAGIMQSGAMVPSDAVDGVYGNEIFDLLASDAGCGSASDKLACLRGVSSDT SWALL: NTYKGKPLFRAGIMQSGAMVPSDAVDGIYGNEIFDLLASNAGCGSASDKLACLRGVSSDT EP0200 LEDATNNTPGFLAYSSLRLSYLPRPDGVNITDDMFALVREGKYASVPVIIGDQNDEGTFF SWALL: LEDATNITPGFLAYSSLRLSYLPRPDGVNITDDMYALVREGKYANIPVIIGDQNDEGTFF EP0200 GTSSLNVTTDAEARQYFTQSFVHASDAELDTLMTAYPQDITQGSPFDTGVLNALTPQFKR SWALL: GTSSLNVTTDAQAREYFKQSFVHASDAEIDTLMTAYPGDITQGSPFDTGILNALTPQFKR EP0200 ISAVLGDLAFIHARRYFLNHYTGGTKYSFLSKQLSGLPVLGTFHSNDIVFQDYLLGSGSL SWALL: ISAVLGDLGFTLARRYFLNHYTGGTKYSFLSKQLSGLPVLGTFHSNDIVFQDYLLGSGSL EP0200 IYNNAFIAFATDLDPNTAGLLVKWPEYTSSSQSGNNLMMINALGLYTGKDNSRTAGYDAL SWALL: IYNNAFIAFATDLDPNTAGLLVKWPEYTSSSQSGNNLMMINALGLYTGKDNFRTAGYDAL 

EP0200 FSNPPSFFV ::::::::

SWALL: FSNPPSFFV